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Probe Trajectory Interpolation for 3D Reconstruction of Freehand Ultrasound

Pierrick Coupé^{a,b,d,*}, Pierre Hellier^{a,b,d} Xavier Morandi^{a,b,d,c}
Christian Barillot^{a,b,d}

^a*University of Rennes I, CNRS, IRISA - UMR 6074, Campus de Beaulieu,
F-35042 Rennes, France*

^b*INRIA, VisAGeS U746 Unit/Project, IRISA, Campus de Beaulieu, F-35042
Rennes, France*

^c*University Hospital of Rennes, Department of Neurosurgery, rue H. Le Guillou,
F-35043 Rennes, France*

^d*INSERM, VisAGeS U746 Unit/Project, IRISA, Campus de Beaulieu, F-35042
Rennes, France*

Abstract

Three-dimensional (3D) Freehand ultrasound uses the acquisition of non parallel B-scans localized in 3D by a tracking system (optic, mechanical or magnetic). Using the positions of the irregularly spaced B-scans, a regular 3D lattice volume can be reconstructed, to which conventional 3D computer vision algorithms (registration and segmentation) can be applied. This paper presents a new 3D reconstruction method which explicitly accounts for the probe trajectory. Experiments were conducted on phantom and intra-operative datasets using various probe motion types and varied slice-to-slice B-scan distances. Results suggest that this technique improves on classical methods at the expense of computational time.

1 Introduction

Ultrasonography has become a very popular medical imaging modality thanks to its low cost, real time image formation capability and non invasive nature. Due to its many attributes, ultrasound has been used in neurosurgery for the last two decades [1]. Several studies demonstrated that ultrasonography can

* Corresponding author.

Email address: pcoupe@irisa.fr (Pierrick Coupé).

be used in the location of tumors, definition of their margins, differentiation of internal characteristics and detection of brain shift and residual tumoral tissue [2].

Despite its advantages, the lack of 3D information in traditional 2D ultrasound imaging prevents reproductivity of examinations, longitudinal follow-up and precise quantitative measurements. To overcome these limits and produce a 3D representation of the scanned organs, several techniques exist: mechanically-swept acquisitions, freehand imaging [3], mechanical built-in probes and 2D phased-array probes [4]. The two first approaches are based on the reconstruction of a 3D regular lattice from 2D B-scans and their positions, whereas 3D probes directly acquire 3D images. The main advantages of freehand imaging, compared to other 3D approaches, are flexibility, low cost and large organs examination capabilities. Moreover, compared to 3D probes, the image quality and the field of view are better suited to clinical applications [5, 6].

Freehand imaging techniques consist of tracking a standard 2D probe by using a 3D localizer (magnetic, mechanical or optic). The tracking system continuously measures the 3D position and orientation of the probe. This 3D position is used for the localization of B-scans in the coordinate system of the localizer. In order to establish the transformation between the B-scan coordinates and the 3D position and orientation of the probe, a calibration procedure is necessary [7, 8]. Calibration is needed to estimate the transformation matrix linking the different coordinate systems (spatial calibration), but also the latency between image and position time stamps (temporal calibration). The localization accuracy of B-scan pixels in the 3D referential system depends on the calibration procedure. A review of calibration techniques is presented in [9].

To analyze the sequences of B-scans, two types of approaches can be used: the reslicing (without reconstruction) or the true 3D reconstruction including interpolation step. The first is used by the StradX system [10] and enables the analysis of the data without reconstruction. The sequence of B-scans can be arbitrarily resliced and distance/volume measurements are performed without reconstruction. This strategy is very powerful for manual analysis of 3D datasets. However, 3D isotropic reconstruction is still necessary in the clinical context when automatic segmentation or registration procedures are required. The second approach is based on the interpolation of the information within the B-scans to fill a regular 3D lattice thus creating a volumetric reconstruction. Due to the non uniform distribution of the B-scans, this step is acutely expensive with respect to computation time and reconstruction quality: an efficient reconstruction method should not introduce geometrical artifacts, degrade nor distort the images. To resolve this problem several methods were proposed. The most common ones are Pixel Nearest-Neighbor (PNN) [11], Voxel Nearest-Neighbor (VNN) [10, 12] and Distance-Weighted interpolation

(DW) [13, 14].

Due to its simplicity of implementation and its reduced computation time, the most straightforward reconstruction algorithm is the PNN method. This algorithm is divided into two stages: the bin-filling and the hole-filling [15]. The bin-filling stage consist in searching, for each pixel in every B-scan. The nearest voxel which is filled with the value of the pixel. Secondly, the remaining gaps in the 3D voxel array are filled *via* a hole-filling method. Usually, the hole-filling method is a local average of filled voxels. Although the PNN method is fast and simple to implement, this approach generates artifacts. Contrary to the PNN method, the VNN approach does not require the hole-filling stage because all voxels are filled in one step using the value of the nearest pixel obtained by orthogonal projection on the nearest B-scan. In the DW interpolation approach, each voxel is filled with the weighted average of pixels from the nearest B-scans (see section 2.1 and Fig. 1 for a detailed explanation). The set of pixels or interpolation kernel is defined either by a spherical neighborhood [13], or by projection on the two nearest B-scans [14]. Then, all pixel intensities of this set are weighted by the inverse distance to the voxel to calculate voxel intensity. A complete survey of these three methods is presented in [16]. These approaches are designed to reduce computation time, at the cost of a lower reconstruction quality compared to more elaborated methods.

More elaborated methods were recently developed in order to increase the reconstruction quality. Firstly, the registration based approach consists in reconstructing a 3D volume after a non-rigid registration of each successive B-scans. In [17], a linear interpolation between the two nearest pixels is used to calculate the intensity voxel. This technique is notably used to avoid artifacts due to tissue motion. Some studies focus on the improvement of the interpolation step using radial basis functions (RBF) [16], weighted Gaussian convolution [6, 18] or Rayleigh model for intensity distribution [19]. Finally, the optimization of the axis orientation and voxel size are discussed in [6]. Nevertheless, the quality improvement obtained with these approaches induces computational burden.

The paper is organized as follows. Section 2 describes the proposed reconstruction method using the 3D probe trajectory (PT) information. Section 3 briefly describes the evaluation framework and compares the proposed method with VNN and DW methods. Finally, in section 4 the advantages and limitations of the PT method are discussed and further improvements are outlined.

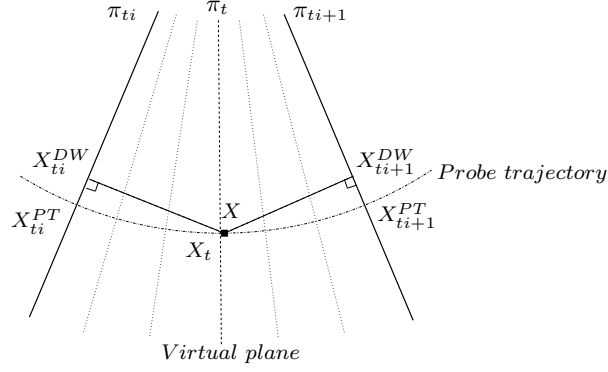


Fig. 1. Illustration of DW and PT principles. The two orthogonal projections for DW interpolation method and the construction of a “virtual” plane π_t containing X for PT method.

2 Method

This work builds on the distance weighted interpolation and proposes to incorporate probe trajectory information. The distance weighted interpolation is first presented in section (2.1). Then, the probe trajectory information is incorporated in section (2.2).

2.1 Distance Weighted Interpolation (DW)

At each point X of the reconstructed volume, the linear interpolation amounts to computing:

$$f_n(X) = \frac{1}{G} \sum_{ti \in K_n(X)} g_{ti} \tilde{f}(X_{ti}) \quad (1)$$

where K_n is the interpolation kernel. In other words, K_n is the set of the different indexes of the B-scans that are involved in the interpolation, n is the interpolation order. For a given interpolation degree, the n closest B-scans before X and the n closest B-scans after X are considered. For the DW interpolation, X_{ti} is the orthogonal projection of X on the ti^{th} B-scan. $\tilde{f}(X_{ti})$ is the intensity at position X_{ti} and is obtained by bilinear interpolation. Finally, G is the normalization constant with $G = \sum g_{ti}$, where g_{ti} is the distance between X and X_{ti} (see Fig. 1).

2.2 Probe Trajectory Interpolation (PT)

The orthogonal projection of points to the nearest B-scans is a straightforward solution. However, it does not take into account the relationship between a given point and its projections. As seen in section 1, registration based in-

interpolation uses homologous points to interpolate, thus increasing the computational burden. We propose to incorporate the probe trajectory into the interpolation process. In other words, homologous points are defined as being successive points along the probe trajectory.

We believe there is correlation between the underlying signal structure and the probe trajectory. When imaging cross-sections of a tubular structure for instance, the intuitive displacement of the probe that follows the Structure Of Interest (SOI) will lead to some correlation between the probe trajectory and the anatomical structure (see the cerebral falx Fig. 11). In intra-operative exams, we observed that the surgeon was concentrated in keeping the focus of the US scans on the SOI (i.e. the lesion). This is confirmed by observing the location of the SOI, which is kept at the same position in the x-y place during the sequence (see Fig. 2). Therefore, we think that the introduction of probe

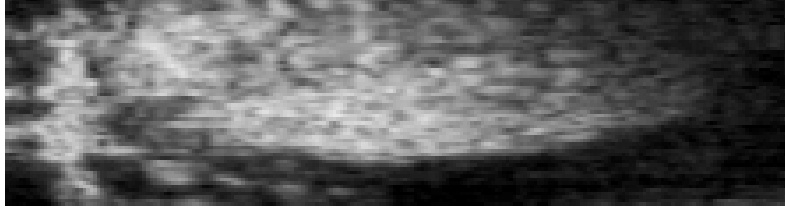


Fig. 2. A longitudinal reslice of the non reconstructed intra-operative data (i.e. the stack of B-scans). The x position in B-scans (horizontal axis of the reslice) of the structure of interest is correlated along the sequence, the vertical axis of the reslice corresponding to the B-scans latencies. The cerebral falx is visible at left and the lesion at center.

trajectory into the interpolation process is relevant.

Instead of using orthogonal projections as in classical DW, we propose to project along the probe trajectory. Firstly, the time stamp $t \in \mathbb{R}$, $t \in [t_i, t_{i+1}]$ of the “virtual plane” π_t is estimated. The “virtual plane” is the plane which passes through X in the sense of the probe trajectory (see Fig 1). Then, t is used to compute the “virtual plane” parameters (translation and rotation) by interpolation of π_{t_i} and $\pi_{t_{i+1}}$ positions. Finally, the 2D coordinates of X_t (the projection of X on π_t) are used to obtain the projections of X on π_{t_i} and $\pi_{t_{i+1}}$ in the sense of the probe trajectory.

2.2.1 Determination of the “virtual” plane time stamp

Under the assumption that the probe motion is constant between two consecutive B-scans, the latency ratio is equal to the distance ratio:

$$t = \frac{d_{ti+1}}{d_{ti} + d_{ti+1}}(t_i) + \frac{d_{ti}}{d_{ti} + d_{ti+1}}(t_i + 1) \quad (2)$$

where d_{ti} is the distance (in the sense of orthogonal projection) between the current voxel and the B-scan of time stamp ti ($d_{ti} = \|X - X_{ti}^{DW}\|$). The assumption of constant probe speed between two slices is justified by the frame rate. The lowest frame rate is usually 10Hz, which means that 100ms separate two frames. It is therefore reasonable to assume a constant motion magnitude between two frames (i.e. no significant acceleration).

Once the time stamp of the “virtual” plane is computed, the probe position can be interpolated.

2.2.2 Determination of the “virtual” plane parameters

The position of each B-scan is defined by 3 translations and 3 rotations. Thus the interpolation of origin position and rotation parameters is needed. We use the Key interpolation for the translations and the Spherical Linear Interpolation (SLERP) for the rotations.

2.2.2.1 Interpolation of origin position For the origin of the B-scan, a cubic interpolation is used to estimate the origin of the “virtual” plane at time stamp t . The Key function is used to carry out a direct cubic interpolation and is defined as:

$$\varphi(t) = \begin{cases} (a+2)|t|^3 - (a+3)t^2 + 1 & \text{if } 0 \leq |t| < 1, \\ a|t|^3 - 5at^2 + 8a|t| - 4a & \text{if } 1 \leq |t| < 2, \\ 0 & \text{if } 2 \leq |t| \end{cases} \quad (3)$$

With $a = -\frac{1}{2}$, φ is a C^1 function and a third order interpolation is obtained [20]. In practice, four B-scans are used for cubic interpolation. This seems to be an optimal trade-off between computational time and reconstruction quality. For example, the interpolation of the origin position along x axis T_x reads as:

$$T_x(t) = \sum_{k=t_i-1}^{t_i+2} T_x(k) \varphi(t-k) \quad (4)$$

2.2.2.2 Interpolation of rotation parameters The rotation parameters of each B-scan are converted into a quaternion which is a compact representation of rotations within a hyper-complex number of rank 4:

$$q = w + ix + jy + kz \quad (5)$$

This representation of rotations, allows to take into account the coupling of rotations during the interpolation step. The quaternion representing the rotations of the “virtual” plane is obtained through a Spherical Linear Interpolation (SLERP) [21] at time stamp t :

$$q_t = q_{ti} \frac{\sin((1-t)\theta)}{\sin\theta} + q_{ti+1} \frac{\sin(t\theta)}{\sin\theta} \quad (6)$$

where q_{ti} and q_{ti+1} are the unit quaternions corresponding to B-scans of time stamps ti and $ti+1$; and θ represents the angle between q_{ti} and q_{ti+1} computed as:

$$\theta = \cos^{-1}(q_{ti} \cdot q_{ti+1}) \quad (7)$$

The orientation of the “virtual” plane is contained in q_t . Then, X_{ti}^{PT} and X_{ti+1}^{PT} are obtained directly, since they have the same 2D coordinates (defined in each B-scans) as X_t .

2.3 Labeling

It is possible that a part of the reconstructed volume is visible on several time stamps (or view points) of the B-scans sequence. These different time stamps are computed during the labeling step so as to track this information and fully exploit the speckle decorrelation. In this way, a label vector L_X containing time stamps of the nearest B-scans is built for each voxel. In case of a simple translation, $L_X = (t_1, t_1 + 1)$ is the label vector of voxel X while t_1 and $t_1 + 1$ are the time stamp of the nearest planes. For more complex probe motions with multiple scanning angles, L_X is composed of several couples of time-consecutive B-scans: $L_X = ((t_1, t_1 + 1), (t_2, t_2 + 1), \dots)$. Afterward, L_X is used to build K_n which also depends on interpolation degree n , thus $K_n = ((t_1 - n + 1, \dots, t_1 + n), (t_2 - n + 1, \dots, t_2 + n), \dots)$. For instance, with an interpolation degree equals to 1 and two view points (see Fig. 3), the “virtual” plane between (π_{t1}, π_{t1+1}) is first computed to estimate the distance of X to (π_{t1}, π_{t1+1}) . Then, the “virtual” plane between (π_{t2}, π_{t2+1}) is computed to estimate the distance of X to (π_{t2}, π_{t2+1}) . Indeed, all the positions of the nearest B-scans are not used at the same time to estimate the probe trajectory, only the consecutive B-scans in time are used simultaneously. Nonetheless, all the pixels on the nearest B-scans are used simultaneously to estimated the intensity of voxel X .

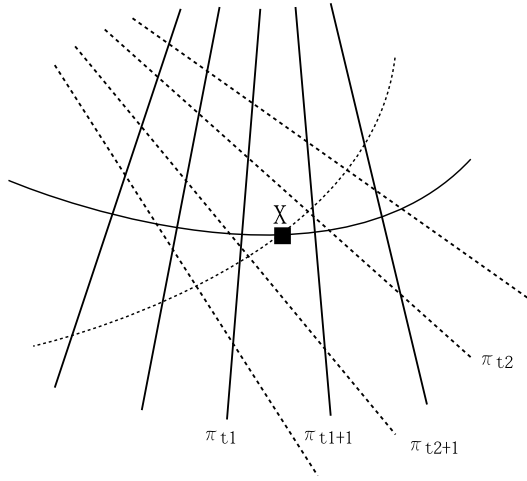


Fig. 3. Illustration of a multi-scanned voxel X at two different time stamps t_1 and t_2 . For an interpolation degree equals to 1, the two couples of B-scans (π_{t1}, π_{t1+1}) and (π_{t2}, π_{t2+1}) are taken into account to evaluate the intensity of X (i.e $K_n = ((t_1, t_1 + 1), (t_2, t_2 + 1))$)

3 Material

3.1 Ultrasound Phantom sequences

A Sonosite system with a cranial 7 – 4MHz probe was used to acquire the ultrasound images. The positions of the B-scans was given by a magnetic miniBIRD system (Ascension Technology Corporation) mounted on the US probe. The StradX software [10] was used to acquire images and 3D position data. The phantom is a CIRS Inc.¹ 3D ultrasound calibration phantom containing two calibrated volumetric ellipsoids test objects. At the acquisition depth, only one of the ellipsoids is visible in the field of view. The two sequences used for the experiments are composed of 510×441 B-scans (204 B-scans for fan motion and 222 B-scans for translation motion, see Fig. 4).

3.2 Ultrasound Intra-operative sequences

For intra-operative sequences the sinuosity cranial probe was coupled with the Sononav Medtronic system in an image-guided neurosurgery context. Contrary to the miniBIRD tracker, the Sononav system is based on an optical tracking to estimate the spatial probe positions sent to the neuronavigation system. The sequences were acquired during neurosurgical procedures after the craniotomy

¹ <http://www.cirsinc.com>

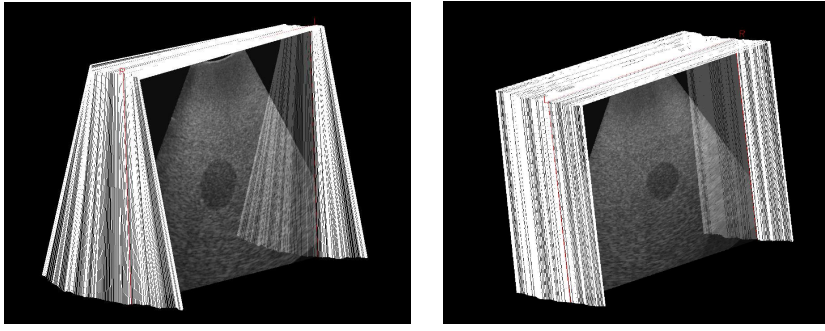


Fig. 4. B-scans sequences used during evaluation. Left: fan sequence. Right: translation sequence.

step but before opening the dura. US-sequence1 is composed of 59 B-scans (223×405) and US-sequence2 of 46 B-scans (223×405), see Fig. 5.

3.3 Magnetic Resonance Intra-operative sequences

In freehand ultrasound imaging the validation step is not an easy task because the “ground truth” does not exist. In order to overcome this problem, magnetic resonance sequences were built on geometry of the ultrasonic intra-operative sequences (see Fig. 5 at bottom). Firstly, the intra operative trajectories were used to re-slice the preoperative magnetic resonance volume of the patient. Then, a stack of MR-scans was built on the images obtained by the re-slicing. Finally, the reconstructed MR volume was compared to the “ground truth” (i.e the corresponding preoperative MRI volume). As the US-sequences, the MR-sequence1 is composed of 59 MR-scans (223×405) and MR-sequence2 of 46 MR-scans (223×405), see Fig. 5.

The following evaluation aims at studying the impact of the probe trajectory incorporation independently of the compounding, the sequences follow simple fan and/or translation motion.

4 Evaluation framework

The performance of the proposed method was compared with two other interpolation approaches: the VNN technique used in StackX [10] and the DW method presented in [14]. For the VNN method, each voxel is projected on the nearest B-scan and its luminance interpolated bilinearly. In the DW technique, each voxel is projected orthogonally on the $2n$ nearest B-scans and its luminance is interpolated (see section 2.1 and Fig. 1).

Within this evaluation framework, the influence of two parameters was stud-

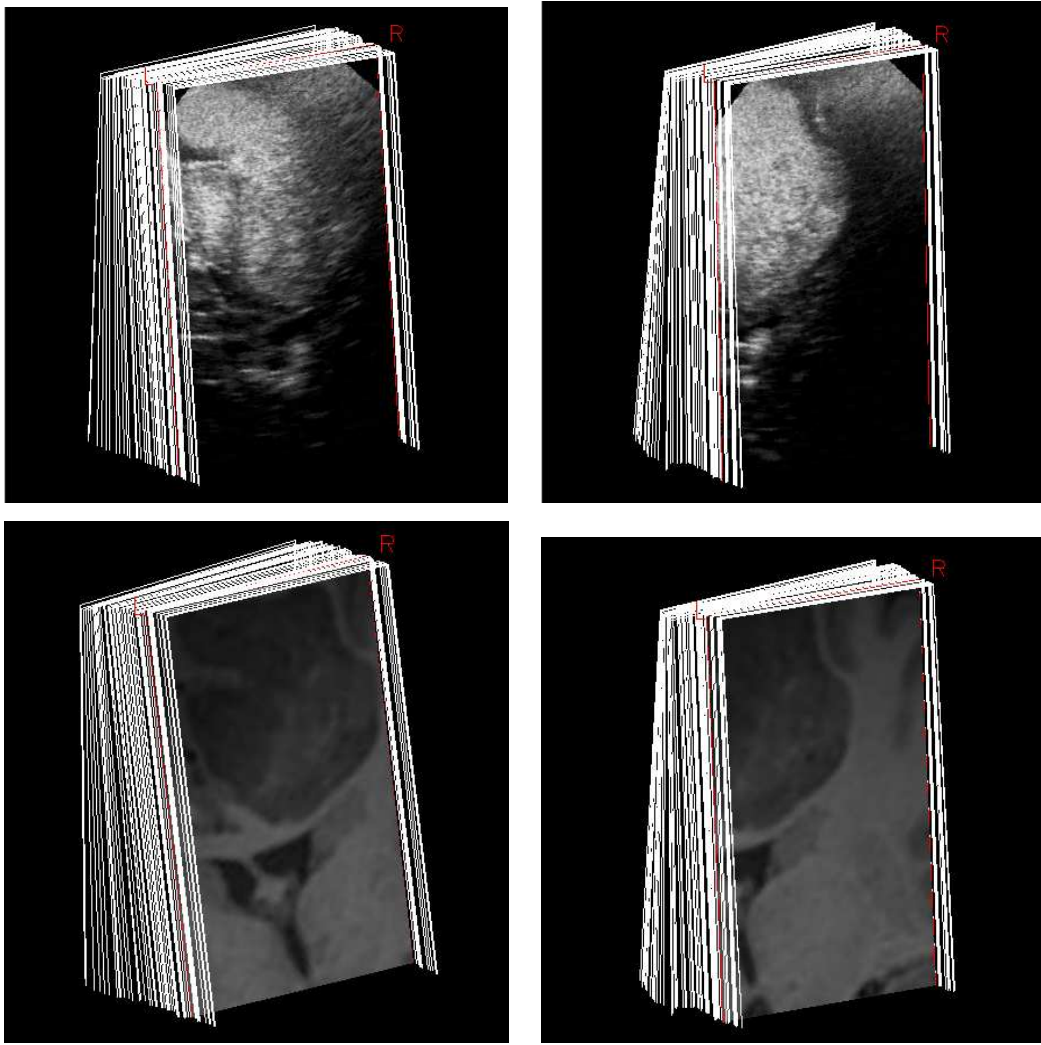


Fig. 5. Top: Intra-operative B-scans sequences of brain used during evaluation. The low-grade glioma and ventricles are visible in white. Left: US-sequence1. Right: US-sequence2. Bottom: Intra-operative MR-scans sequences of brain built on B-scans sequences. Left: MR-sequence1. Right: MR-sequence2. In MR images the low-grade glioma and the ventricles appears in gray.

ied:

- “The distance between two consecutive B-scans”. The sequence is sub-sampled thanks to SelectSX², which simulates a lower frame acquisition rate, in order to artificially increase the distance between 2 consecutive B-scans. The evaluation framework studies the probe trajectory interpolation impact according to the distance between two consecutive B-scans.
- “The size of the interpolation kernel”. For ultrasound phantom sequences, the removed B-scans are reconstructed with different methods and different interpolation degrees (from 1 to 2 for DW and PT methods).

² <http://mi.eng.cam.ac.uk/~rwp/stradx/utilities.html>

4.1 For Ultrasound Phantom Data

To assess the reconstruction quality, evaluation data can be created from any image sequence: given a sequence of 3D freehand US, each B-scan is removed from the sequence, and then reconstructed by using the other B-scans. This “leave one out” procedure is performed for each B-scan. The Mean Square Error (MSE) is used as quality criterion:

$$MSE(t) = \frac{1}{P} \sum_{j=1}^P (\tilde{I}_t(x_j) - I_t(x_j))^2 \quad (8)$$

where I_t is the original image at time t (removed from the sequence), \tilde{I}_t the reconstructed image and P is the number of pixel in this B-scan. After MSE estimation for all B-scans of the sequence, we compute the mean μ and the standard deviation σ of the reconstruction error.

$$\mu = \frac{1}{N - 2n} \sum_{t=n+1}^{N-n} MSE(t) \quad (9)$$

$$\sigma^2 = \frac{1}{N - 2n} \sum_{t=n+1}^{N-n} (MSE(t) - \mu)^2 \quad (10)$$

N is the total number of B-scan in the sequence and n the interpolation kernel degree.

4.2 For Magnetic Resonance Sequences

Contrary to ultrasound phantom data, for MR-sequences the “ground truth” is known. The evaluation framework directly compares the reconstructed volume \tilde{V} and the corresponding volume V in preoperative MRI. Firstly, the MR sequence, obtained by reslicing the pre-operative MRI accordingly to the US trajectory, was reconstructed with the three evaluated methods. Secondly, the corresponding MR volume V (in terms of field of view and resolution) was computed using cubic interpolation. Finally, the reconstructed volumes obtained with VNN, DW and PT were compared to the “ground truth” V (see Fig. 6). The MSE between V and \tilde{V} , obtained with the different methods, was evaluated after removing the background (i.e. voxels zero intensity). The transformation matrix given by the neuronavigator is used to register the US and MR images. This registration is not corrected with an external registration procedure since only the intra-operative trajectory is required and used to create the stack of MR-scans. This means that tracking and calibration errors are removed : therefore reconstruction errors can be studied independently of other sources of errors.

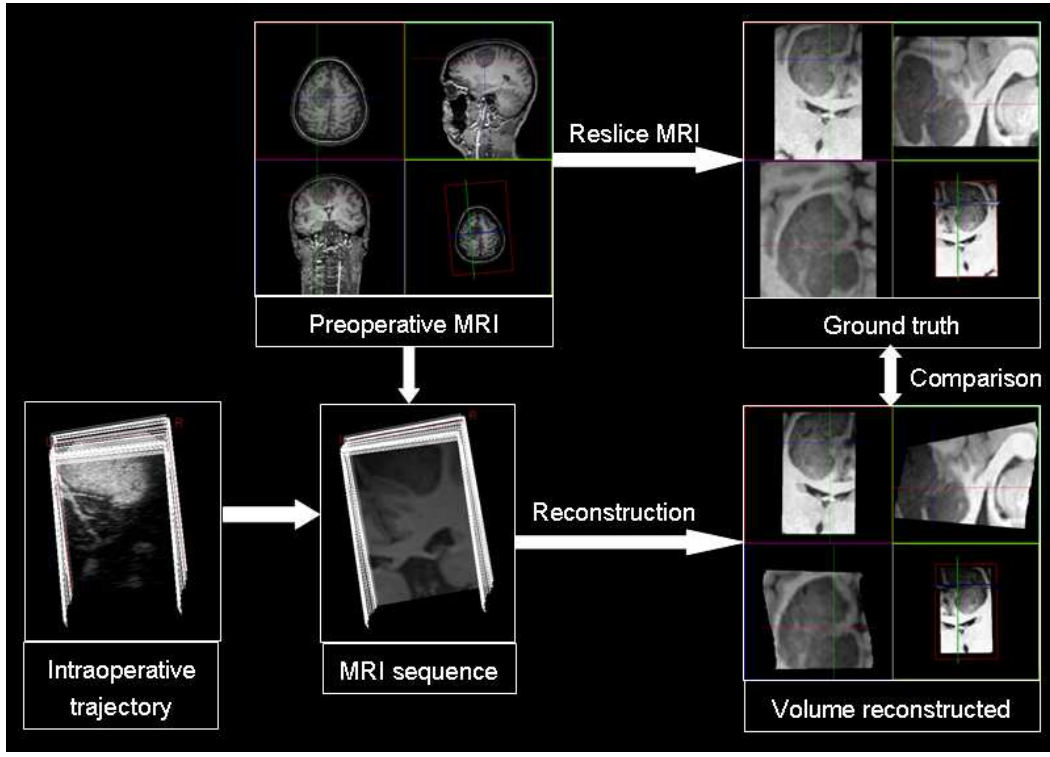


Fig. 6. Illustration of the validation framework used for the MR-sequences.

5 Results

The reconstructions were performed on a P4 3.2 Ghz with 2Go RAM. In order to only compare interpolation times, the total computational time is divided as follows: L_{time} corresponds to labeling step and I_{time} corresponds to interpolation time. The labeling step consists in constructing the label vector K_n for each voxel X (described in 2.3).

5.1 Interpolation Function

In this section, the Key interpolation of rotation parameters is compared with the theoretically optimal approach described in paragraph 2.2.2.2. The direct interpolation of each parameter can be considered as an approximation of the Spherical Linear Interpolation (SLERP) because the coupling of rotations is not taken into account. However, for small rotations, this approximation leads to very similar results. Table 1 compares the reconstruction quality using the SLERP and interpolation approximation. The SLERP interpolation is theoretically correct, but experiments showed that reconstruction results are similar. This is due to the continuity of probe motion and the proximity of two consecutive B-scans. Considering these results, the Key interpolation will be

used in the rest of the paper in order to decrease the computational burden.

			SLERP			KEY		
Motion	Mean Distance	L_{time}	μ	σ	I_{time}	μ	σ	I_{time}
Fan	0.3 mm	414s	21.5	10.9	355s	21.5	11.0	114s
	0.9 mm	145s	47.9	16.1	370s	47.8	15.9	111s
Translation	0.2 mm	42s	16.3	2.9	440s	16.2	2.9	138s
	0.6 mm	20s	45.1	12.1	413s	45.1	12.1	138s

Table 1

Error measures composed of mean μ and standard deviation σ for the Spherical Linear Interpolation (SLERP) and the Key interpolation. Results on phantom sequence indicate that the improvement in terms of interpolation quality with SLERP was not significant whereas the computational time increased.

5.2 Phantom sequence

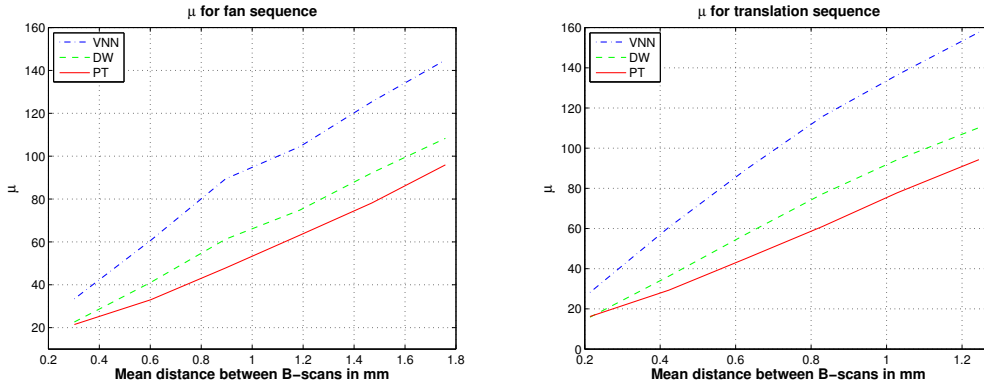


Fig. 7. Variation of error reconstruction relatively to distance between two consecutive B-scans on phantom sequence with interpolation degree equals to 1. Left fan motion, right translation motion. Three methods are evaluated: VNN, DW and PT. The PT method outperforms others methods especially on sparse data.

Results on phantom sequences (described in section 3) are presented in Figure 7 and Table 2. Figure 7 shows the influence of the mean distance between two B-scans on the reconstruction error with two types of motion (i.e. translation and fan). Table 2 presents the error and computation time for different interpolation degrees (1 and 2). In all cases, the PT method outperforms the VNN and the DW methods especially on sequences with few B-scans. The probe trajectory is especially relevant to compensate for the sparsity of data. When the distance of a given point to the B-scans considered in the interpolation increases, the orthogonal projections and probe trajectory differ significantly. This distance introduces artifacts for the DW method. For distances close to $0.2mm$, DW and PT methods are equivalent. The difference between PT

Motion	Distance between B-scans	Interpolation degree	VNN			DW			PT		
			μ	σ	I_{time}	μ	σ	I_{time}	μ	σ	I_{time}
Fan	$L_{time} = 414s$	$n = 1$	33.4	13.5	20s	22.6	11.6	44s	21.5	11.0	114s
		$n = 2$				26.2	12.5	46s	22.9	10.6	122s
	$L_{time} = 215s$	$n = 1$	60.7	21.6	21s	41.1	16.9	31s	33.0	14.2	114s
		$n = 2$				46.6	17.6	44s	38.2	14.1	124s
	$L_{time} = 145s$	$n = 1$	89.4	25.2	20s	61.3	19.4	30s	47.8	15.9	111s
		$n = 2$				67.6	18.6	43s	56.2	14.2	118s
Translation	$L_{time} = 42s$	$n = 1$	28.2	9.6	20s	15.9	4.8	37s	16.2	2.9	138s
		$n = 2$				19.1	6.4	49s	18.3	3.5	149s
	$L_{time} = 27s$	$n = 1$	60.6	22.9	21s	36.3	13.9	32s	29.3	8.6	138s
		$n = 2$				43.0	14.5	47s	35.8	8.7	147s
	$L_{time} = 20s$	$n = 1$	89.3	29.6	20s	57.1	18.3	31s	45.1	12.1	138s
		$n = 2$				63.7	17.5	46s	52.9	11.1	146s

Table 2

Error measures composed of mean square error μ and standard deviation σ for the different methods (see section 4). L_{time} is the time spent for labeling, while I_{time} is the time spent for interpolation. Results indicate that the PT method obtains better results than the VNN and the DW methods. The improvement in terms of reconstruction quality is obtained at the expense of a slight computational increase.

and DW, in terms of mean square error for translation and fan motion was expected to be greater. We think that the phantom images do not contain enough structures to really show the reconstruction quality improvement in case of fan motion. Apart from the ellipsoid in the image center, the image contains only speckle. For the fan sequence, the difference between PT and DW, in terms of projection, is substantial for voxels that are far from the rotation center, that is to say in deep areas. The deep regions do not convey structural information but mostly speckle.

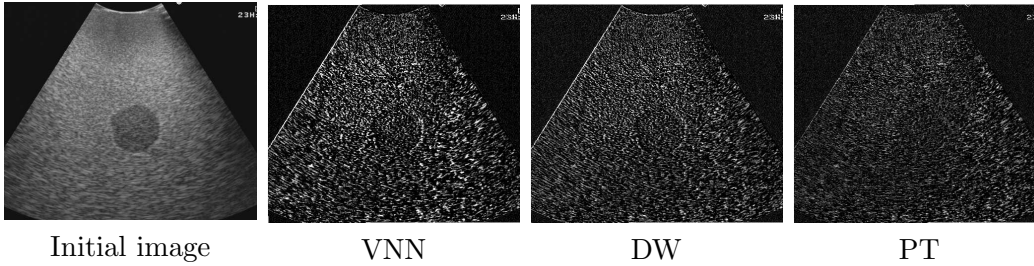


Fig. 8. Differences between original (left) and reconstructed B-scan for fan sequence with $n = 1$. From left to right: the Voxel Nearest Neighbor, the Distance Weighted interpolation and the Probe Trajectory methods. This shows that the error between the reconstructed B-scan and the initial image is visually lower with the PT method, especially on the contours of the object.

Figure 8 shows the differences between the original and the reconstructed B-scans with the VNN, DW and PT methods. Visually, the PT reconstruction appears closer to the original B-scan, what fits the numerical results of Table

5.3 MR Intra-operative sequences

Figure 9 shows the influence of the mean distance between two consecutive B-scans on the reconstruction error. The mean square error is computed between the “Ground Truth” and the reconstructed volume. The PT method outperforms the VNN and DW approaches especially on sparse data. Figure 10 presents slices extracted from initial MR volume and the reconstructed MR volume. Visually, the PT method preserves more the edges and the contrast. Compared to DW method, the PT method improves the reconstruction result especially on edges whose direction is correlated with probe trajectory.

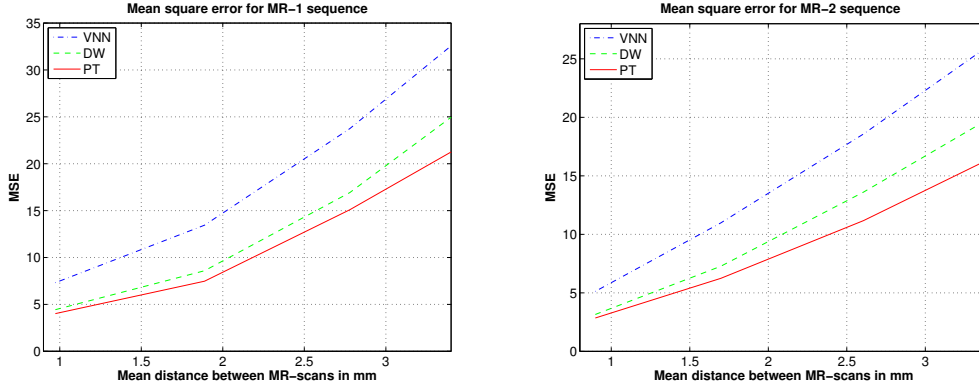


Fig. 9. Variation of mean reconstruction error relatively to the distance between two consecutive MR-scans with interpolation degree of 1. Left MR-sequence1, right MR-sequence2. Three methods are evaluated: VNN, DW and PT. The PT method outperforms others methods especially on sparse data.

5.4 US Intra-operative sequences

The dimension of the reconstructed volume are $263 \times 447 \times 306$ for US-sequence1 and $286 \times 447 \times 234$ for US-sequence2 with resolutions of $(0.188, 0.172, 0.180)$. The reconstruction process, with a multithreading implementation on an Intel Pentium Dual Core CPU at 3.40GHz with 2Go RAM, takes 220s for US-sequence1 and 154s for US-sequence2.

The reconstructions of B-scans dataset US-sequence1 are presented in Figure 11 and US-sequence2 are presented in Figure 12. Visually, the VNN method leads to many discontinuities and creates artificial boundaries (see image at the top right of Fig. 12). The DW method generally smooths out the edges and spoils the native texture pattern of US image more than PT (see at the bottom of Fig. 12).

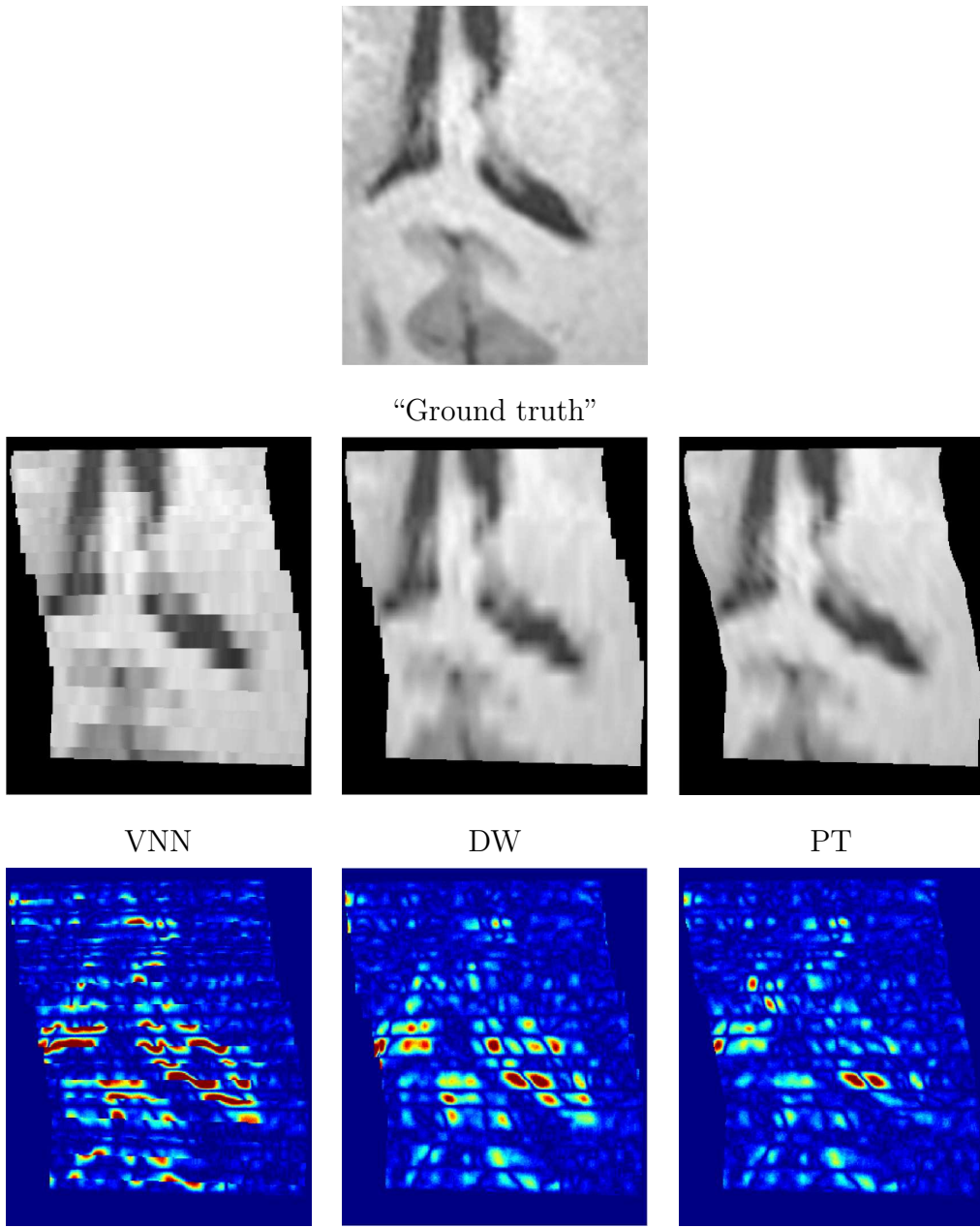
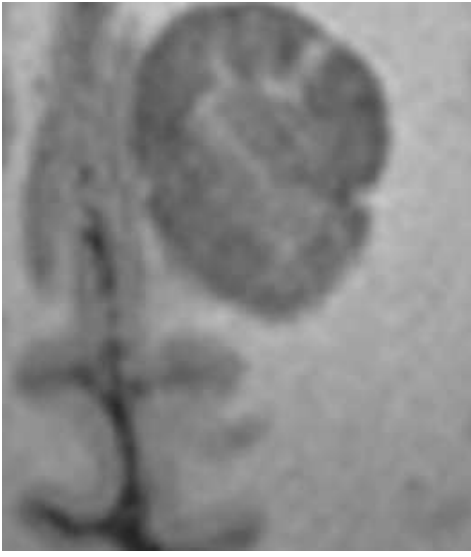
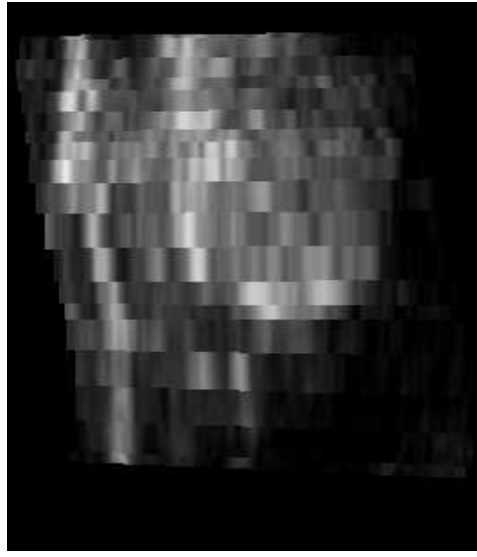


Fig. 10. Results for MR-sequences of brain. Top: the “ground truth” and the reconstructions obtained *via* the different methods for a mean distance between MR-scans of $1.8mm$. On the bottom: the images of the difference between the “ground truth” and the reconstructed volume. Visually, the PT method preserves more the edge continuity and the contrast. The difference images show that the PT method creates less artifacts and appears closer to the “ground truth”, especially on ventricles edges.

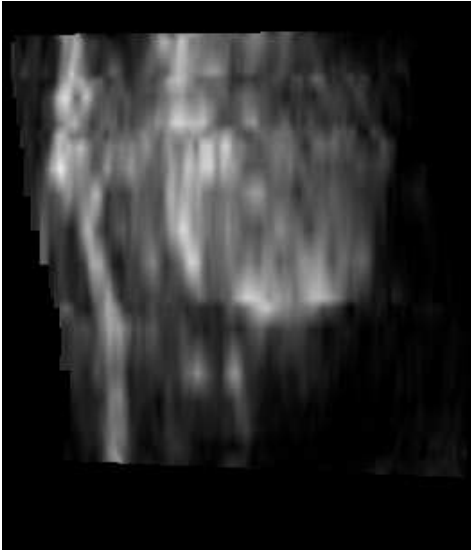
The actual implementation of methods can be largely optimized as compared to StackSX, which produces all reconstruction processes of translation phantom sequence in 10 seconds with the VNN method (62s with our implementation of VNN). Although the labeling step needs significant improvement, this



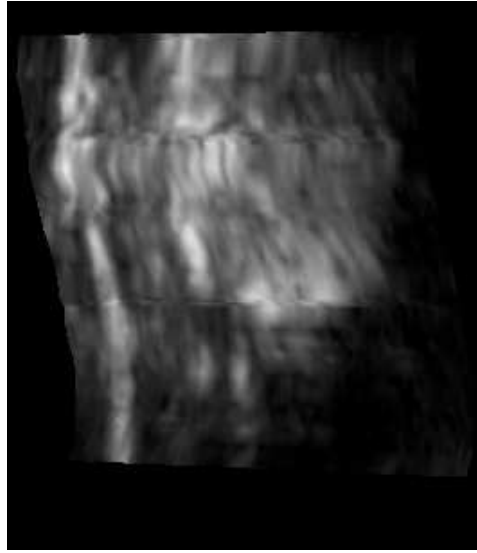
MRI



VNN



DW



PT

Fig. 11. Results for US-sequence1 of brain. Top: the preoperative MRI and the US reconstruction obtained with the VNN method. On the bottom: the reconstruction obtained with DW and PT approaches. The low-grade glioma and the cerebral falx appear in gray in MR image and in white in US images. From left to right the VNN, DW and PT methods. The images highlight the inherent artifacts of VNN (i.e. discontinuities) and DW (i.e. blur) methods. These results underline that the PT method preserves the edges continuity and the native texture of US image more than VNN and DW method.

study aims to compare computation time between identical implementations of methods. The increased quality of reconstruction for the PT method is obtained at the expense of a slight increase of the computation time. Nonetheless this side effect seems to be reasonable with regards to the reconstruction qual-

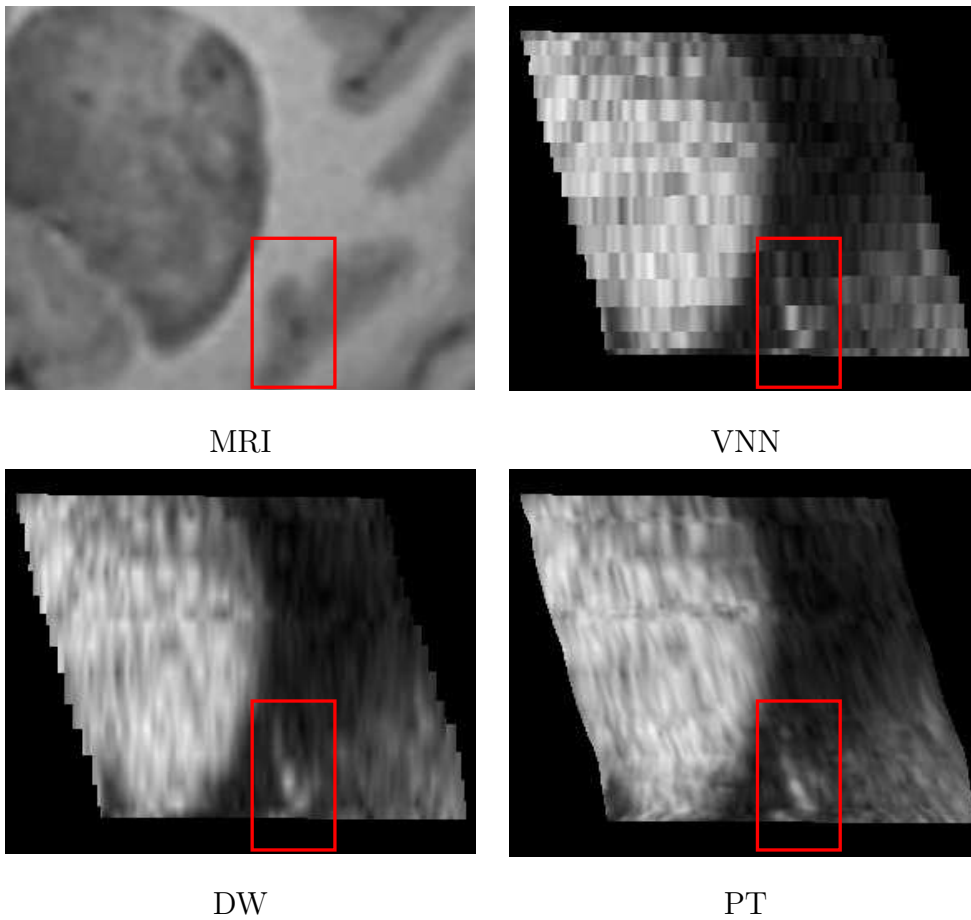


Fig. 12. Results for US-sequence2 of brain. Top: the preoperative MRI and the US reconstruction obtained with the VNN method. On the bottom: the reconstruction obtained with DW and PT approaches. The low-grade glioma appears in gray in MR image and in white in US images. Visually, the PT method preserves more the edges continuity especially on sulci edges (see at the center bottom of images).

ity. Contrary to more elaborated techniques like non-rigid registration or RBF, which are considerably computationally expensive, the PT approach offers an attractive compromise between computation time and reconstruction quality.

6 Discussion and conclusion

This paper presented a 3D freehand ultrasound reconstruction method explicitly taking into account the probe trajectory information. Through an evaluation framework, it shows that the proposed method performs better than traditional reconstruction approaches (i.e. Voxel Nearest Neighbor and Distance Weighted interpolation) with a reasonable increase of the computation time. The main limitation of PT method is the assumption of constant probe speed between two slices. Nonetheless, this hypothesis is reasonable when using a

decent frame rate (more than 10Hz). Moreover, the direct interpolation of rotation parameters instead of Spherical Linear Interpolation (SLERP) does not introduce artifacts but accelerates the reconstruction process. The evaluation results underline the relevance of probe trajectory information, especially on sequences with low frame rate acquisitions or large distances between two consecutive B-scans. The PT method is a trade-off between reconstruction error and computational time. The results show that the hypothesis of a correlation between the signal structure and the probe trajectory is relevant. Indeed, in practice, the probe trajectory and signal structure are correlated because the medical practitioner tends to follow the structure of interest (see on Fig. 11). The precise localization of anatomy and pathology within the complex 3D geometry of the brain remains one of the major difficulties of neurosurgery. Thus, image-guided neurosurgery (IGNS) is an adequate context to experiment the proposed method because the reconstruction quality is of utmost importance and the time dedicated to image reconstruction is limited. The PT method is thus interesting for applications where an accurate reconstruction is needed in a reasonable time. Further work should be pursued for comparing the PT reconstruction approach with registration based approaches [17]. Registration based approaches avoid artifacts of slice misregistration due to errors in tracking data (i.e. trajectory) and/or tissue deformation. The compensation of tissue motion can be especially interesting in neurosurgery due to the complex problem of soft tissue deformations also known as brainshift. However, since the registration is a computationally expensive process, the benefit for image reconstruction should be studied. Then, our implementation could be largely optimized using graphic library implementation (ex: *OpenGL*) or grid computation especially for IGNS purpose. Finally, the impact of PT reconstruction on registration (mono and multimodal) to compensate for errors of localization and brainshift needs to be investigated further. Indeed, the current pitfalls of the neuronavigator system are: the errors caused by geometrical distortion in the preoperative images, registration, tracking errors [22], and brainshift.

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